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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

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A16K 37/62, C07H 17/00, 15/12 A61K 31/70	A1	13) International Publication Date: 21 March 1991 (21.03.91
(22) International Filing Date: 5 June 199 (30) Priority data: 401,613 31 August 1989 (31.08. (60) Parent Application or Grant (61) Related by Continuation	.89) 01,613 (C 89 (31.08	(75) Inventors/Applicants (for US only): ROSSI, John, J. [US. US]; 346 Cimmeron Trail, Glendora, CA 91740 (US) CHANG, Pairoj [US/US]; 949 Avenida Loma Vists San Dimas, CA 91773 (US). KAPLAN, Bruce, E. [US. US]; 825 N. Indian Hill, Claremont, CA 91711 (US). (74) Agent: IRONS, Edward, S.; 919 18th Street, N.W., Suit 800, Washington, DC 20006 (US). (81) Designated States: AU, CA, DE*, FR (European patent GB, IT (European patent), JP, US.

(54) Title: CHIMERIC DNA-RNA CATALYTIC SEQUENCES

DRDRD-1

GGUGCGAGAGCGUCAGUAUUAAGCGG CCACGCTCTCGCA TCATAATTCGCC - HIV 792-817 AGC AGG =RNA G(A)T C C G GT

(57) Abstract

This invention provides chimeric DNA/RNA catalytic molecules useful to cleave RNA sequences. The invention specifically provides two different chimeric DNA-RNA-DNA-RNA-DNA catalytic molecules which are targeted to cleave HIV-1 RNA sequences. These chimeric molecules include DNA sequences which flank a catalytic RNA center. interaction with the HIV-1 substrate RNAs is achieved by Watson-Crick base pairing of the DNA flanking sequences with HIV-1 RNA. The catalytic ribonucleotide center cleaves the phosphodiester bond of the substrute HIV-! RNA at the expected location.

^{*} See back of page

Summary of the Invention

This invention provides chimeric DNA/RNA catalytic molecules useful to cleave RNA sequences. The invention specifically provides two different chimeric DNA-RNA-DNA-RNA-DNA catalytic molecules which are targeted to cleave HIV-1 RNA sequences. These chimeric molecules include DNA sequences which flank a catalytic RNA center. Interaction with the HIV-1 substrate RNAs is achieved by Watson-Crick base pairing of the DNA flanking sequences with HIV-1 RNA. The catalytic ribonucleotide center cleaves the phosphodiester bond of the substrate HIV-1 RNA at the expected location.

General Description of the Invention

In general the catalytic molecules of the invention function as hammerhead or hairpin ribozymes. The preferred molecular construct consists of two known RNA catalytic sequences each flanked by a DNA sequence at the respective 3' and 5' termini and coupled by a DNA sequence at the corresponding 5' and 3' termini. These molecules may accordingly be represented by the formulae I and II::

I. 3' X - AAAG - Y - AGUAGUC - Z 5'

or

II. 3' X - CAAAG - Y - AGUAGUC - Z 5' in which X, Y and Z are DNA sequences and AAAG, CAAAG and AGUAGUC are catalytic RNA sequences.

The flanking X and Z components may be any DNA sequences that allow base pairing with the substrate RNA at appropriate positions adjacent to the substrate cleavage site. These flanking sequences may be phosphodiester, phosphorothicate, methyl phosphonate, methyl phosphorate or similar moieties.

Y may be any DNA sequence that base pairs <u>inter</u> se in the manner required for catalytic cleavage of

130 × F.

the substrate by the RNA sequences preferably as shown in base paired form in Formula III:

The catalytic molecules of this invention can be synthesized in known manner by commercially available DNA synthesizers such as those produced by Applied Biosystems or Milligen. See, e.g., Perreault, et al, supra.

The X and Z sequences may be substituted at the respective 3' and 5' ends with ligands to facilitate cell entry, targeting within the cell and ultimate stability of the catalysts. Such ligands include by way of example but not of limitation: other nuclotides, proteins, carbohydrates, lipids, steroid hormones and cholesterol.

The catalytic molecules of the invention are administered by known and available delivery agents or systems, including, but not limited to, liposomes, defective viral particles, viral capids, and standard DNA/RNA transfective procedures.

Description of the Figures

Figure 1 illustrates one catalytic molecule of the invention base paired to an HIV-1 sequence. The RNA portion of the molecule is encircled.

Figure 2 illustrates a second catalytic molecule of the invention base paired to another HIV-1 sequence. The RNA portion of the molecule is encircled.

Figure 3A depicts a ribonuclease A digestion of the catalytic molecule of Figure 1 as compared with an equivalent all DNA molecule. The conditions were - 12

10 units of commercial (Sigma) pancreatic ribonuclease in 2XSSC buffer added to the oligonucleotides which were in 10 microliters of 50 mM Tric-HCl buffer (pH 8.0). The RNAse was incubated with the sample for 10 minutes before the 32-p end labelled DRDRD or DNA molecules were electrophoresed in a 15t polyacrylamide gel containing 8M urea. The gel was autoradiographed for 10 minutes to get the exposure depicted.

Figure 3B depicts a cleavage reaction involving the catalytic molecule of Figure 1 under conditions described in Chang, et al., Clinical Biotechnology, 2:23-31 (1990).

EXAMPLE I

The catalytic molecule of Figure 1 was synthesized in known manner utilizing an automated oligonucleotide synthesizer manufactured by Applied Biosystems, Inc.

The result of ribonuclease A digestion of the catalytic molecule is shown by Figure 3A.

The catalytic molecule produced, as described, was used to cleave each of a 610 nuleotide long (S-610) and a 170 nucleotide long HIV-1 gag transcript. In brief, the buffer was 50 mM Tris-HC1, pH 7.5, lmM EDTA, 10mM MgCl₂ at approximately 1 pmole of target, 3 pmole of ribozyme or DNA. The reactions were carried out at 37°C. for 12 hours. The substrate was either a 610 nucleotide long HIV-1 gag containing transcript (S-610) or a 172 nucleotide long HIV-1 gag containing transcript (S-172). The 5' cleavage product is indicated for both.

In Figure 3B the 5' cleavage product is shown for both transcripts. The 3' cleavage product for the 610 target is not visible due to poor reproduction of

Specific cleavage of an HIV-1 5' LTR splice site with a similar catalytic molecule has also been obtained.

CLAIMS

- 1. A catalytic molecule capable of cleaving an HIV-1 RNA sequence at a known ribozyme cleavage site said molecule having the formula
 - 3' X AAAG Y AGUAAGUC Z 5'

or

3' X - CAAAG - Y - AGUAAGUC - Z 5' in which X and Z are DNA sequences that base pair with an RNA substrate at positions juxtaposed to said known cleavage site,

AAAG, CAAAG and AGUAGUC are RNA sequences,
Y is a DNA sequence that base pairs <u>inter</u> se in a

manner required to permit said RNA sequences to cleave said substrate at said cleavage site.

- 2. The catalytic molecule shown by Figure 1.
- 3. The catalytic molecule shown by Figure 2.
- 4. A catalytic molecule, as defined by Claim 1, in which said RNA sequence is an HIV-1 sequence.
- 5. A catalytic molecule, as defined by Claim 4, in which said HIV-1 sequence is the HIV-1 sequence shown by Figure 1.
- 6. A catalytic molecule, as defined by Claim 4, in which the HIV-1 sequence is the HIV-1 sequence shown by Figure 2.
- 7. A catalytic molecule capable of cleaving an RNA sequence, said molecule having catalytic RNA moieties linked to first and second DNA moieties which base pair with the substrate RNA sequences flanking the cleavage site and interconnected by a third DNA sequence which base pairs <u>inter</u> se to facilitate said cleavage.

FIG. 1 DRDRD-1

5' GGUGCGAGAGCGUCAGUAUUAAGCGG 3' - HIV 792-817
CCACGCTCTCGCA) TCATAATTCGCC 5' - HIV 792-817

A C UG
A G
G C
G C
G C
A G
G T

FIG. 2 DRDRD #2

2/2



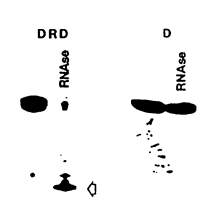
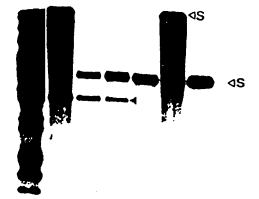


FIG. 3B



	INTERNATIONAL SE	ARCH REPORT		
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III. DOCU	IMENTS CONSIDERED TO BE RELEVANT 14			
Calegory *	Citation of Document, 14 with Indication, where approp	riale, of the relevant passages 17	Relevant to Claim No. 14	
A,P	Chemical Abstract, Volume 112, No. 7, issued 1 - 7 12 February 1990 (Columbus, Chio, U.S.A.) W. Gerlach, et al, "Synthetic Ribozymes for in Vivo Inactivation of Prokaryotic or Eukaryotic RNA Transcripts", See pages 336-337, column 2, See the abstract No. 51284j, Eur. Pat. Appl. EP 321,201 21 June 1989.			
A,P	Chemical Abstract, Volume 112, No. 19, issued 07 May 1990 (Columbus, Ohio, U.S.A.) N. Sarver, et al, "Ribozymes as Potential Anti-HIV-1 Therapeutic Agents", See page 420, column 2, See the abstract No. 17548q, Science, 1990, 247 (4947), 1222-5 (Eng).			
A,P	Chemical Abstract, Volume 112, No. 7, issued 1 - 7 12 February 1990 (Columbus, Ohio, U.S.A.), M. Cotten, et al, "Ribozyme Mediated Destruction of RNA in Vivo", See page 501, column 1, See the abstract No. 52942j, EMBO J, 1990, 8(12), 3861-6 (Eng).			
"A" do	cial categories of cited documents: 13 ocument defining the general state of the art which is not onsidered to be of particular relevance artier document but published on or after the international ling date ocument which may throw doubts on priority claim(s) or hich is cited to establish the publication date of another listion or other special reason (as specified) ocument referring to an oral disclosure, use, exhibition or ther means ocument published prior to the international filing date but liter than the priority date claimed	"T" later document published alte or priority date and not in co-cited to understand the principal content of particular relevantion cannot be considered novel involve an inventive step "T" document of particular relevannot be considered to involve and the constitution being the art. "4" document member of the same of Maillian of this laterations.	nflict with the application but iple or theory underlying the ance: the claimed invention or-cannot be considered to ance: the claimed invention re an inventive step when the ne or more other such docu- ing obvious to a person skilled the patent lamily	
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111 00000	international Application No.	PCT/US90/03102
Category .	ENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHE	(T)
Caragod)	Citation of Document, 44 with Indication, where appropriate, of the relevant passages 17	Relevant to Claim No
A,P	Nature, volume 344, issued 05 April 1990, J. Peneault, et al., Mixed Decryribo — and Ribooligoracleotides with Catalytic activity see pages 565-567.	1-7
A,P	Proceeding of the National Academy of Sciences, Volume 86, no. 23, issued December 1989 (U.S.A.) F.H. Cameron, et al., 'Specific Gene Suppression by Engineered Ribozymes in Monkey Cells', see pages 9139 - 9143.	
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TORTHER INFORMA	TION CONTINUED FROM THE SECOND SHEET	PC170590703102	
et al. Restric Manufac	al Abstracts, Volume 110, No. 21, issued 1989, (Columbus, Ohio, U.S.A.) T. R. Cech , "RNA Ribozyme Polymerases, Dephosphorylases, tion Endoribonucleases and Methods for Their cture", See page 226, column 2, See the ct No. 187321K, PCT Int. Appl. W08804,300 e 1988.	1 - 7	
V. OBSERVATION	S WHERE CERTAIN CLAIMS WERE FOUND UNSEARCHABLE!	<u> </u>	
This international search	report has not been established in respect of certain claims under Article 17(2) (a) is		
1. Claim numbers	, because they relate to subject matter t not required to be searched by this Aut	ar the following reasons:	.
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VI. OBSERVATIONS	WHERE UNITY OF INVENTION IS LACKING?		2.5
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